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## (54) Title: METHODS AND MATERIALS RELATING TO DNA BINDING PROTEINS

## (57) Abstract

DNA sequences associated with regulation of early stages of cell growth are described. Illustratively provided are human and mouse origin DNA sequence encoding early growth regulatory ("Egr") proteins which include "zinc finger" regions of the type involved in DNA binding. Immunological methods and materials for detection of Egr proteins and hybridization methods and materials for detection and quantification of Egr protein related nucleic acids are also disclosed.

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"METHODS AND MATERIALS RELATING TO  
DNA BINDING PROTEINS"

5

CROSS-REFERENCE TO RELATED APPLICATIONS

10        This is a continuation-in-part application of  
co-pending U.S. Patent Application Serial No. 179,587,  
filed April 8, 1988.

15

BACKGROUND OF THE INVENTION

20        The present invention relates generally to DNA  
binding regulatory proteins and more particularly to DNA  
sequences encoding early growth regulatory proteins  
possessing histidine-cysteine "zinc finger" DNA binding  
25        domains, to the polypeptide products of recombinant  
expression of these DNA sequences, to peptides and poly-  
peptides whose sequences are based on amino acid  
30        sequences deduced from these DNA sequences, to anti-  
bodies specific for such proteins and peptides, and to  
procedures for detection and quantification of such pro-  
teins and nucleic acids related thereto.

35

      Among the most significant aspects of mam-  
malian cell physiology yet to be elucidated is the  
precise manner in which growth factors (e.g., hormones,  
neurotransmitters and various developmental and  
differentiation factors) operate to effect the regula-  
tion of cell growth. The interaction of certain growth  
factors with surface receptors of resting cells appears  
to rapidly induce a cascade of biochemical events  
thought to result in nuclear activation of specific  
growth related genes, followed by ordered expression of  
other genes. Analysis of sequential activation and  
expression of genes during the transition from a resting  
state ("G<sub>0</sub>") to the initial growing state ("G<sub>1</sub>") has  
been the subject of substantial research. See, gener-

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ally, Lau et al., Proc. Nat'l. Acad. Sci. (USA), 84, 1182-1186 (1987). Much of this research has involved 5 analysis of the expression of known genes encoding suspected regulatory proteins (such as the proto-oncogenes, c-fos and c-myc) following mitogen stimulation. An 10 alternative approach has involved attempts to identify genes activated by mitogenic stimuli through differential screening of cDNA libraries prepared from resting 15 cells following exposure to serum and specific growth factors. See, e.g., Lau et al., EMBO Journal, 4, 3145-3151 (1985). See also, Cochran et al., Cell, 33, 939-947 (1983), relating to the cloning of gene 20 sequences apparently regulated by platelet derived growth factor.

Of interest to the background of the invention 25 is the continuously expanding body of knowledge regarding structural components involved in the binding of regulatory proteins to DNA. Illustratively, the so-called receptor proteins are believed to bind to DNA by 30 means of zinc ion stabilized secondary structural fingers premised on folding of continuous amino acid sequences showing high degrees of conservation of 35 cysteines and histidines and hydrophobic residues. See, e.g., Gehring, TIBS, 12, 399-402 (1987). For example, a "zinc finger" domain or motif, present in Xenopus transcription factor IIIA (TF IIIA), as well as the Drosophila Kruppel gene product and various yeast proteins, involves "repeats" of about 30 amino acid residues wherein pairs of cysteine and histidine residues are coordinated around a central zinc ion and are thought to form finger-like structures which make contact with DNA. The histidine-cysteine (or "CC-HH") zinc finger motif, as opposed to a cysteine-cysteine ("CC-CC") motif of steroid receptors, is reducible to a consensus sequence represented as C-X<sub>2-4</sub>-C-X<sub>3</sub>-F-X<sub>5</sub>-L-X<sub>2</sub>-H-X<sub>3</sub>-H wherein C represents cysteine, H represents

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histidine, F represents phenylalanine, L represents leucine and X represents any amino acid. [See, Klug et al. TIBS, 12, 464-469 (1987); Blumberg et al., Nature, 328, 443-445 (1987); and Schuh et al., Cell, 47, 1025-1032 (1986).]

Of particular interest to the background of the invention is the recent report of Chowdhury et al., Cell, 48, 771-778 (1987), relating to an asserted "family" of genes encoding proteins having histidine-cysteine finger structures. These genes, designated "mkrl" and "mkrl2", appear to be the first such isolated from mammalian tissue and are not correlated to any early growth regulatory events.

There continues to exist a need in the art for information concerning the primary structural conformation of early growth regulatory proteins, especially DNA binding proteins, such as might be provided by knowledge of human and other mammalian DNA sequences encoding the same. Availability of such DNA sequences would make possible the application of recombinant methods to the large scale production of the proteins in prokaryotic and eukaryotic host cells, as well as DNA-DNA and DNA-RNA hybridization procedures for the detection, quantification and/or isolation of nucleic acids associated with these and related proteins. Possession of such DNA-binding proteins and/or knowledge of the amino acid sequences of the same would allow, in turn, the development of monoclonal and polyclonal antibodies thereto (including antibodies to protein fragments or synthetic peptides modeled thereon) for use in immunological methods for the detection and quantification of early growth regulatory proteins in fluid and tissue samples as well as for tissue specific delivery of substances such as labels and therapeutic agents to cells expressing the proteins. In addition, DNA probes based on the DNA sequences for these mammalian early growth

5 regulatory proteins may be of use in detecting gene markers used for the diagnosis of those clinical disorders which are linked to the marker genes.

BRIEF SUMMARY OF THE INVENTION

10 The present invention provides novel purified and isolated DNA sequences encoding mammalian early growth regulatory ("Egr") proteins which comprise one or 15 more histidine-cysteine zinc finger amino acid sequences putatively providing DNA binding (and hence DNA replication or transcription regulatory) capacity. In 20 presently preferred forms, novel DNA sequences of the invention comprise genomic and cDNA sequences encoding human and mouse early growth regulatory proteins. 25 Alternate DNA forms, such as "manufactured" DNA, prepared by partial or total chemical synthesis from nucleotides, are also within the contemplation of the invention.

30 Operative association of Egr-encoding DNA sequences provided by the invention with homologous or heterologous species expression control DNA sequences, 35 such as promoters, operators, regulators and the like, allows for in vivo and in vitro transcription to form messenger RNA which, in turn, is susceptible to translation to provide Egr proteins in large quantities. In one presently preferred DNA expression system practiced according to the invention, Egr-encoding DNA is operatively associated with a bacteriophage T3 or T7 RNA promoter DNA sequence allowing for in vitro transcription and translation in a cell free system. Incorporation of novel DNA sequences of the invention into prokaryotic and eucaryotic host cells by standard transformation and transfection processes involving suitable viral and circular DNA plasmid vectors is also within the contemplation of the invention and is expected to

provide useful proteins in quantities heretofore unavailable from natural sources. Illustratively, fragments of DNA encoding Egr protein of the invention have been incorporated in plasmid vectors resulting in expression by transformed E.coli hosts of fusion proteins sharing immunological characteristics of Egr protein. Use of mammalian host cells is expected to provide for such post-translational modifications (e.g., truncation, glycosylation, and tyrosine, serine or threonine phosphorylation) as may be needed to confer optimal biological activity on recombinant expression products of the invention.

Also provided by the present invention are novel, presumptively mitogen sensitive, DNA sequences involved in regulation of the transcription of Egr-encoding DNA, which sequences are expected to have utility in the efficient recombinant expression of Egr proteins as well as proteins encoded by other structural genes. In addition, the DNA sequences may be used as probes to detect the presence or absence of gene markers used for the diagnosis of clinical disorders linked to those gene markers.

Novel polypeptide products of the invention include polypeptides having the primary structural conformation (i.e., amino acid sequence) of Egr proteins or fragments thereof, as well as synthetic peptides, analogs thereof, assembled to be partially or wholly duplicative of amino acid sequences extant in Egr proteins. Proteins, protein fragments, and synthetic peptides of the invention are expected to have therapeutic, diagnostic, and prognostic uses and also to provide the basis for preparation of monoclonal and polyclonal antibodies specifically immunoreactive with Egr proteins, as well as to provide the basis for the production of drugs for use as competitive inhibitors or potentiators of Egr. Preferred protein fragments and

5 synthetic peptides of the invention include those  
duplicating regions of Egr proteins which are not  
involved in DNA binding functions (i.e., regions other  
than the zinc fingers). Most preferred are peptides  
10 which share at least one continuous or discontinuous  
antigenic epitope with naturally occurring Egr pro-  
teins.

15 Antibodies of the invention preferably bind  
with high immunospecificity to Egr proteins, fragments,  
and peptides, preferably recognizing epitopes which are  
not common to other proteins, especially other DNA bind-  
ing proteins.

20 Also provided by the present invention are  
novel procedures for the detection and/or quantification  
of Egr proteins and nucleic acids (e.g., DNA and mRNA)  
25 specifically associated therewith. Illustratively, anti-  
bodies of the invention may be employed in known immuno-  
logical procedures for quantitative detection of Egr  
proteins in fluid and tissue samples. Similarly, DNA  
30 sequences of the invention (particularly those having  
limited homology to other DNAs encoding DNA binding  
proteins) may be suitably labelled and employed for the  
35 quantitative detection of mRNA encoding the proteins.  
Information concerning levels of Egr mRNA may provide  
valuable insights into growth characteristics of  
cells.

Among the multiple aspects of the present  
invention, therefore, is the provision of (a) novel  
purified and isolated Egr-encoding DNA sequences set out  
in Figures 1A, 3, and 4 as well as (b) Egr-encoding DNA  
sequences which hybridize thereto under hybridization  
conditions of the stringency equal to or greater than  
the conditions described herein and employed in the  
initial isolation of DNAs of the invention, and (c)  
synthetic or partially synthetic DNA sequences encoding  
the same, or allelic variant, or analog Egr polypeptides

which employ, at least in part, degenerate codons. Correspondingly provided are viral or circular plasmid 5 DNA vectors incorporating such DNA sequences and pro- caryotic and eucaryotic host cells transformed or trans- fected with such DNA sequences and vectors as well as 10 novel methods for the recombinant production of Egr proteins through cultured growth of such hosts and iso- lation from the hosts or their culture media.

Preferred polypeptide products of the inven- 15 tion include those wholly or partially duplicating the deduced sequence of the amino acid residues set out in Figures 1A and 3 (i.e., mouse "Egr-1" and human 20 "EGR2"). Other preferred polypeptides include fusion proteins such as cro- $\beta$ -galactosidase/Egr-1 and bovine growth hormone/Egr-1.

Presently preferred antibodies of the inven- 25 tion include those raised against synthetic peptides partially duplicating deduced Egr amino acid sequences 30 of Figures 1A and 3 (e.g., the synthetic peptides H-L-R- Q-K-D-K-K-A-D-K-S-C, the first 12 amino acid residues of which duplicate mouse Egr-1 residues 416-427 with the 35 last cysteine added for coupling to KLH; and C-G-R-K-F- A-R-S-D-E-R-K-R-H-T-K-I duplicating mouse Egr-1 residues 399-415). The antisera against the first peptide is designated VPS10 and comprises a preferred antibody of the invention.

As employed herein, the term "early growth regulatory protein" shall mean and include a mammalian DNA binding protein encoded by DNA whose transcription temporally corresponds to cellular events attending the G<sub>0</sub>/G<sub>1</sub> growth phase transition. As employed herein, "histidine-cysteine zinc finger amino acid sequence" shall mean and include the following sequence of amino acids C-X<sub>2-4</sub>-C-X<sub>3</sub>-F-X<sub>5</sub>-L-X<sub>2</sub>-H-X<sub>3</sub>-H wherein C represents cysteine, H represents histidine, F represents phenylalanine, L represents lysine, and X represents an amino acid.

5 Other aspects and advantages of the present invention will be apparent upon consideration of the following detailed description thereof which includes numerous illustrative examples of the practice of the invention, reference being made to the drawing wherein:

10 Figure 1A provides a 3086 base nucleotide sequence for a mouse Egr-1 DNA clone as well as a deduced sequence of 533 amino acid residues for the protein; Figure 1B provides a partial restriction map of 15 Egr-1 DNA clones together with information concerning the position of the protein coding sequence and the locus of amino acids providing for histidine-cysteine 20 zinc fingers;

25 Figure 2 provides an amino acid sequence alignment of the DNA binding domain of mouse Egr-1 in comparison with a zinc finger consensus sequence, with the *Drosophila Kruppel* sequence and with the "finger 2" sequence of *Xenopus TFIIIA* protein;

30 Figure 3 provides a 2820 base nucleotide sequence for a human EGR2 cDNA clone as well as a deduced sequence of 456 amino acids for the protein;

35 Figure 4 provides a 1200 base nucleotide sequence of a mouse Egr-1 genomic clone, specifically illustrating the 5' non-transcribed regulatory region thereof comprising bases -935 through +1; and

Figure 5 provides a restriction map and organization of the mouse Egr-1 genomic clone mgEgr-1.1 and a comparison to mouse Egr-1 cDNA.

#### DETAILED DESCRIPTION

The following examples illustrate practice of the invention. Example 1 relates to the preparation and structural analysis of cDNA for mouse Egr-1. Example 2 relates to confirmation of the presence of an Egr DNA sequence on human chromosome 5. Example 3 relates to the in vitro transcription and translation of mouse

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5       Egr-1 cDNA. Example 4 relates to production of anti-  
bodies according to the invention. Example 5 relates to  
the isolation and characterization of genomic DNA which  
encodes mouse Egr-1. Example 6 relates to the isolation  
and characterization of cDNA encoding human EGR2.  
10      Example 7 relates to preparation, in an E. coli host, of  
a recombinant fusion protein including a portion of the  
deduced amino acid sequence of mouse Egr-1. Example 8  
relates to use of DNA probes of the invention in the  
15      quantitative detection of EGR1 mRNA.

20      These examples are for illustrative purposes  
only and are not intended in any way to limit the scope  
of the invention.

#### EXAMPLE 1

25      Preparation and Structural  
Analysis of cDNA for Mouse Egr-1

30      Isolation of DNA encoding a mammalian early  
growth regulatory protein including one or more histi-  
dine-cysteine zinc finger amino acid sequences was per-  
formed substantially according to the procedures des-  
cribed in Sukhatme et al., Oncogene Research, 1, 343-355  
35      (1987), the disclosures of which are specifically incor-  
porated by reference herein.

Balb/c 3T3 cells (clone A31) from the American  
Type Culture Collection were grown to confluence in  
Dulbecco's Modified Eagle's medium (DME) supplemented  
with 10% fetal calf serum (FCS). The cells were ren-  
dered quiescent by reduction of the serum concentration  
to 0.75% for 48 hours. To induce the cells from quies-  
cence into growth phase G<sub>1</sub>, the medium was changed to  
20% FCS with cycloheximide added to a final concentra-  
tion of 10 µg/ml.

RNA was extracted from Balb/c 3T3 cells har-  
vested three hours after induction of quiescent cells by

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20% FCS and 10  $\mu$ g/ml cycloheximide. A  $\lambda$ gt10 cDNA library was constructed from this mRNA according to the procedures of Huynh et al., DNA Cloning, Vol. 1, 49-78 (Glover, D., ed., IRL Press, 1985). This library was screened differentially with single stranded cDNA prepared from quiescent cells and from cells exposed to serum and cycloheximide for 3 hours. These  $^{32}$ P-labeled cDNA probes were prepared from poly A<sup>+</sup> RNA as described in St. John, et al., Cell, 16, 443-452 (1979), except that 100  $\mu$ Ci of  $^{32}$ P-dCTP (>3000 Ci/mmol), 0.02 mM cold dCTP and 2-5  $\mu$ g of poly A<sup>+</sup> RNA was used in each reaction. The mean size of the reverse transcribed probes, as assessed by alkaline agarose gel electrophoresis and subsequent autoradiography, was about 700 bases. Replica filter lifts (GeneScreenPlus, NEN-DuPont) were prepared essentially as described by Benton et al., Science, 196, 180-192 (1977), and approximately  $3 \times 10^6$  cpm of  $^{32}$ P-cDNA were used per filter (90 mm diameter). Hybridizations were carried out at 65°C in 1% SDS, 10% dextran sulfate, and 1 M NaCl for a period of 16 hours. The filters were washed twice for twenty minutes each time, first at room temperature in 2 x SSC [Maniatis et al., Molecular Cloning, Cold Spring Harbor Laboratory (New York, 1982)], then at 65°C in 2 x SSC, 1% NaDODSO<sub>4</sub> and finally at 65°C in 0.2 x SSC. Autoradiograms were prepared by exposing the blots for 18 hours at -70°C with an intensifying screen.

A total of 10,000 cDNA clones from the Balb/c 3T3  $\lambda$ gt10 library were differentially screened. Seventy-eight clones were found to hybridize preferentially to single-stranded cDNA from fibroblasts stimulated for 3 hours with 20% FCS and cycloheximide as compared to single-stranded cDNA from quiescent cells. Inserts from these clones were cross-hybridized to each other, resulting in the sorting of forty clones into 7 cDNA families one of which was identified as c-fos.

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Another cDNA clone, referred to as OC68, contained a 2.2 kb insert and was characterized further. This 5 insert was subcloned into the Eco RI site of pUC13 and probes were generated for Northern blot analysis either from the insert or the corresponding pUC plasmid. 10 Figure 1B illustrates a partial restriction digest map of the OC68 clone ("R" representing restriction sites for RsaI) along with that of a shorter clone, OC19t. 15 Two RsaI digestion fragments, derived from the 5' end of clone OC68 and each comprising approximately 130 base pairs, were labeled and employed to re-screen the above-described  $\lambda$ gt10 cDNA library, resulting in the recovery 20 of a 3.1 kb clone, designated OC3.1, shown in figure 1B. This clone was sequenced according to the method of Sanger et al., Proc. Nat'l. Acad. Sci. (USA), 74, 5463 25 (1977). The 3086 base pair sequence obtained is set forth in Figure 1A along with the deduced sequence of 533 amino acid residues for the protein encoded, 30 designated mouse "Egr-1".

The deduced amino acid sequence shows a single long open reading frame with a stop codon (TAA) at position 1858. The most 5', in-frame, ATG, at position 259, 35 is flanked by sequences that fulfill the Kozak criterion (A<sub>n</sub>NN(ATG)G) [Kozak, Nuc. Acids Res., 15, 8125-8131 (1987)]. The sequence region upstream of this ATG is highly GC-rich and results in an absence of in-frame stop codons. The 3' untranslated region (UT) contains two "AT" rich regions (nucleotides 2550-2630 and 2930-2970). Similar sequences are found in the 3' UT regions of several lymphokine and proto-oncogene mRNAs, including granulocyte macrophage colony stimulating factor (GM-CSF), interleukin 1, interleukin 2, interleukin 3 (IL-3),  $\alpha$ ,  $\beta$ , and  $\gamma$  interferons, and c-fos, c-myc, and c-myb [Shaw et al., Cell, 46, 659-667 (1986)]. These sequences may mediate selective mRNA degradation. The presence in the mouse Egr-1 transcript

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of such regions is consistent with its short message half-life. Potential polyadenylation signals (AATAAA) 5 are located at nucleotide positions 1865 and 3066, as well as at position 3053 (AATTAA) [Wickens et al., Science, 226, 1045-1051 (1984)].

10 The deduced amino acid sequence predicts a polypeptide of 533 amino acids with a molecular weight of 56,596. Based on structural considerations, namely a central region containing zinc fingers (described 15 below), the Egr-1 protein can be divided into three domains. The N-terminal portion (amino acid residues 2 to 331) is rich in proline (14.2%) and serine (16%) residues with 7.9% alanines and 7.9% threonines. The C-terminal region (residues 417 to 533) also contains a very high proportion of prolines and serines (15.4 and 25 26.5%, respectively) and 10.3% alanines and 11.1% threonines. The large number of proline residues leads to a secondary structure that probably lacks  $\alpha$ -helices. The central portion of the Egr-1 protein 30 consists of three tandem repeat units of 28-30 amino acids, with the first unit starting at position 332. Each unit conforms almost exactly to the consensus 35 sequence  $TGX_{3F}^YXCX_{2-4}CX_3FX_5LX_2HX_3H$  (see Figure 2), diagnostic of DNA binding zinc fingers [Berg, Science, 232, 485-486 (1986); Brown et al., Nature, 324, 215 (1986); and Brown et al., FEBS Letters, 186, 271-274 (1985)]. Furthermore, the Egr-1 fingers are connected by "H-C links" ( $TGE_{K^R}^P_{Y^F}X$ ) [Schuh et al., Cell, 47, 1025-1032 (1986)] found in the Xenopus TFIIIA gene (between fingers 1, 2, and 3), in the Drosophila Kruppel gap gene [Rosenberg et al., Nature, 319, 336-339 (1986)], and in genes from mouse and Xenopus that cross-hybridize to the Kruppel ( $K^R$ ) finger domains: mkrl, mkr2 [Chowdhury et al., Cell, 48, 771-778 (1987)], and Xfin [Altaba et al., EMBO Journal, 6, 3065-3070 (1987)]. The sequence similarity amongst the Egr-1

5       fingers is 50-70%, whereas the sequence similarity  
between any of the Egr-1 fingers and those present in  
TFIIIA, Kruppel, mkr1, mkr2 or Xfin is 35-40%. Outside  
of the finger domains, it is noteworthy that the Egr-1  
and Kr proteins each contain a very high proportion of  
10      Pro, Ala, and Ser residues [Schuh et al., Cell, 47,  
1025-1032 (1986)]. However, there is no sequence  
similarity in these regions. Thus, Egr-1 and Kr are not  
homologous genes nor is Egr-1 related to mkr1, mkr2,  
15      Xfin, or TFIIIA. The Kr gene contains thirteen copies  
of the hexanucleotide (ACAAAAA), or its complementary  
sequence, eight of which are located within 180 bp  
20      downstream from the Kr TATA box and five are in the 3'  
UT region. These sequences may serve as targets for  
other DNA binding proteins or in Kr gene autoregula-  
25      tion. The Egr-1 cDNA also contains nine copies of the  
ACAAAAA sequence or its complement.

30       Following the work described above, Milbrandt  
[Science, 238, 797-799 (1987)], reported the isolation  
and sequence of a nerve growth factor (NGF) inducible  
cDNA (NGFI-A) from the rat pheochromocytoma PC12 line.  
A comparison of the deduced amino acid sequence of  
35      NGFI-A to that of mouse Egr-1 of Figure 1A reveals 98%  
sequence identity. Thus, mouse Egr-1 and rat NGFI-A are  
homologs. The putative initiation ATG chosen by  
Milbrandt corresponds to position 343 in the Figure 1A  
cDNA sequence, and is 84 nucleotides (28 amino acid  
residues) downstream of the ATG therein designated for  
translation initiation. Both ATG's have a purine at  
position -3 and a G at position +1 and the designation  
represented in Figure 1A of the more 5' ATG as the puta-  
tive start codon is based on the experience of Kozak,  
Nuc. Acids Res., 15, 8125-8131 (1987), even though the  
more 3' ATG is surrounded by the longer Kozak consensus  
sequence (CCG/ACCATGG). Translation of an in vitro  
generated RNA transcript, described infra, selects the  
more 5' ATG for initiation.

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It is noteworthy that a major difference in the deduced sequences of mouse Egr-1 and rat NGFI-A 5 resides in the sequence spanning residues 61-68 of Egr-1 and 33-43 of NGFI-A. The former includes the sequence N-S-S-S-S-T-S-S while the latter includes the sequence 10 N-N-S-S-S-S-S-S-S-S, accounting for the 3 residue difference in length of the putative polypeptides which is not accounted for by the difference in designation of 15 the transcript initiation signal.

15

EXAMPLE 2

20 Human Chromosome Gene Mapping

To determine the human chromosomal localization of the gene corresponding to mouse Egr-1, 25 the OC3.1 and OC19t cDNA clones were hybridized to a panel of rodent x human somatic cell hybrids. Southern blot analysis of the hybrid panel showed concordance between the presence of Egr-1 sequences and human 30 chromosome 5. In situ hybridization to normal human metaphase chromosomes resulted in specific labeling only of chromosome 5, with the largest cluster of grains at 35 5q23-31. Specific labeling of these bands was also observed in hybridizations using an Egr-1 probe which does not contain finger sequences.

This localization is interesting in light of the non-random deletions [del(5q)] in human myeloid disorders (acute myelogenous leukemia) (AML), and myelodysplastic syndromes, that involve this chromosomal region. [Le Beau et al., Science, 231, 984-987 (1986); Dewald et al., Blood, 66, 189-197 (1985); and Van den Berghe et al., Cancer Genet. Cytogenet., 17, 189-255 (1985)]. Fifty percent of patients with therapy related AML show chromosome 5 abnormalities (interstitial deletions or monosomy) and cytogenetic analysis of the deletions has revealed that one segment, consisting of bands

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5q23-31, is absent in the malignant cells of all patients who have aberrations of chromosome 5. These 5 data suggest that loss of a critical DNA sequence leading to hemizygosity (or homozygosity) of a recessive allele may play an important role in the pathogenesis of these disorders, a mechanism substantiated for retinoblastoma. Although genes for a number of growth factors and receptors (IL-3, GM-CSF,  $\beta_2$ -adrenergic receptor, endothelial cell growth factor, CSF-1, c-fms, pDGF 10 receptor) are clustered in or near this region, Egr-1 (by virtue of its zinc fingers) is the only member of this group with potential transcriptional regulatory 15 activity. It is therefore possible that its absence 20 could lead to deregulated cell growth.

EXAMPLE 3

25

In Vitro Expression of Mouse Egr-1 cDNA

30 A 2.1 kb ApaI/ApaI fragment (comprising nucleotides 120-2224 of Figure 1A) was isolated from the OC3.1 DNA clone. This fragment includes the translation start (ATG) codon at nucleotide position 259 designated 35 in Figure 1A. The fragment was blunt-ended with T4 DNA polymerase and cloned into the Bluescript vector KS M13(+) containing a T3/T7 bacteriophage promoter. The (T3) sense transcript was generated and in vitro translated in a standard rabbit reticulocyte lysate system (Promega Biotec, Madison, WI. 53711) including <sup>35</sup>S methionine as a radiolabel. An analogous in vitro transcription system was developed using a BglIII/BglII fragment of OC3.1 (including nucleotides 301-1958 and not including the translation start designated in Figure 1A). The T7 sense transcript was employed in the translation system. Differential characterization of translation products by autoradiographic SDS PAGE indicated that the ATG at nucleotide position 259 is preferred as

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a translation start codon when all potential start sites are present.

5

EXAMPLE 4

10 Preparation of Antibodies:

10 A first synthetic peptide based on the sequence of amino acid residues 416-427 of mouse Egr-1 was prepared and provided with a carboxy terminal cysteine residue. The peptide, H-L-R-Q-K-D-K-K-A-D-K-S-C, was coupled to KLH and employed to immunize New Zealand white rabbits. Animals were initially immunized 20 with 100 µg of the immunogen in Freund's Complete Adjuvant and every two weeks were boosted with 100 µg of immunogen with Freund's Incomplete Adjuvant. Sera, 25 designated VPS10, were isolated after 68 days and displayed an antibody titer of 1:12,800 based on reactivity with the antigen used to prepare the anti- sera.

30

30 A second synthetic peptide, based on residues 399 to 415 of mouse Egr-1, was prepared. The peptide, C-G-R-K-R-A-R-S-D-E-R-K-R-H-T-K-I, was coupled to KLH 35 and used to immunize rabbits as above, resulting in the production of antisera (designated VPS2) with a titer of 1:400.

EXAMPLE 5

Isolation of Genomic Mouse Egr-1 Clone and Characterization of Regulatory Regions

A mouse Balb/c 3T3 genomic library was prepared in a Stratagene (La Jolla, California) vector, λFIX, according to the manufacturer's instructions and probed using 1% SDS, 1 M NaCl, and 10% dextran sulfate at 65°C with stringent final wash in 0.2 x SSC at 65°C with a 2.1 kb ApaI/ApaI fragment and a 3.1 kb Eco RI/Eco

5 RI fragment derived from digestion of pUC13 including the mouse Egr-1 clone OC3.1. One positive clone, from approximately 300,000 screened, was designated mgEgr-1.1 and also hybridized to the extreme 5'-end 120 bp Eco RI-  
Apa I fragment from plasmid OC3.1.

10 A 2.4 kb Pvu-II-PvuII fragment and a 6.6 kb XbaI-XbaI fragment (see Figure 5) derived from the mgEgr-1.1 clone were subcloned into the SmaI and XbaI sites of pUC13 and pUC18 respectively, and the resulting 15 plasmids (designated as p2.4 and p6.6) were used for restriction mapping analysis of transcription initiation sites and for nucleotide sequencing. Marked in Figure 20 4, and listed in Table 1, are possible regulatory elements identified in the 5' flanking sequence of mgEgr-1.1. A putative TATA motif (AAATA) is located 26 nucleotides upstream of the transcription start site. A 25 "CCAAT" type sequence starts at nucleotide -337. Five different regions, each 10 nucleotides in length, located at -110, -342, -358, -374, and -412, are nearly 30 identical to the inner core of the c-fos serum response element (Treisman, R., Cell, 46, 567 (1986)). Each has a 5-6 nucleotide AT rich stretch and is surrounded by 35 the dinucleotide CC on the 5' side and GG on the other. Two potential TPA responsive elements (Lee, W., et al., Cell, 49, 741 (1987) and Angel, P., et al., Cell, 49, 729 (1987)) are located at nucleotides -610 and -867. Four consensus Spl (Briggs, M.R., et al., Science, 234, 47 (1986) binding sequences are at position -285, -649, -700 and -719. In addition, two sequences have been identified that might serve as cAMP response elements (Montimy, M.R., et al., Nature, 328, 175 (1987)) (-138 and -631).

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TABLE 1

Location and Identification of Potential Regulatory Elements			
	<u>Element</u>	<u>Sequence<sup>1</sup></u>	<u>Location<sup>2</sup></u>
10	TATA CCAAT	AAATA CCAAT	-26 to -22 -337 to -333
15	Serum Response Element Consensus		
	GATGTCCATATTAGGACATC CC TA AT GG G C	TCC TTCCATATTAGGGCTTC <u>GTGGCCC-AATATGGCCCTG</u> <u>CAGGCCCTTATATGGAGTGG</u> <u>ACAGACCTTATTGGGCAGC</u> <u>AAACGCCATATAAGGAGCAG</u>	-110 to -91 -342 to -324 -358 to -339 -374 to -355 -412 to -393
20	TPA Responsive Element (AP1 binding site) Consensus		
25	C C TGACT A G A	CTGACTCG CTGACT <u>GG</u>	-610 to -603 -867 to -860
30	Spl binding site	GGGC <sup>GG</sup> GGGC <sup>GG</sup> CCGCC <sup>CC</sup> GGGC <sup>GG</sup>	-285 to -280 -649 to -644 -700 to -695 -719 to -714
35	cAMP Response Element Consensus		
	TGACGTCA	TCACGTCA <u>TGACGGCT</u>	-138 to -131 -631 to -624

1. The underlined bases in the mouse Egr-1 gene sequence are those that do not match the consensus sequence.

2. The location numbers refer to the nucleotides of the mouse Egr-1 gene as indicated in Figure 4.

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To obtain the genomic sequence and the intron-exon gene structure, specific oligonucleotides (17-mers at positions 83, 122, 174, 200, 379, 543, 611, 659, 905, 920, 1000, 1200, 1400, 1600, 1800, 2100, 2353, 2650, 2825) of the OC3.1 cDNA sequence (see Figure 1A) were used as primers for double stranded sequencing of plasmids p2.4 and p6.6. Comparison of the Egr-1 genomic sequence to the Egr-1 cDNA sequence showed the Egr-1 gene consists of 2 exons and a single 700 bp intron (between nucleotide position 556 and 557 as numbered in Figure 1A and as shown in Figure 5). Both the 5' and 3' splice junction sequences (not shown) are in excellent agreement with the consensus boundary sequences. Mount, S.M., Nucleic Acids Res., 10, 459 (1982).

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EXAMPLE 6

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Isolation and Characterization  
of Human EGR2 cDNA

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A human genomic placental library in the vector EMBL3, prepared by Dr. C. Westbrook of the University of Chicago according to procedures described in Frischauff et al., Jour. Mol. Biol., 170, 827-842 (1983), and a human leukocyte cosmid library prepared according to procedures described in Proc. Nat'l. Acad. Sci. (USA), 80, 5225-5229 (1983), were probed with the 2.1 kb ApaI fragment of OC3.1 (described in Example 5) using 1% SDS, 1 M NaCl and 10% dextrose sulfate at 50-55°C with a non-stringent final wash in 2 x SSC at 50-55°C. A single positive clone (designated HG6) was isolated from the first library and four clones (designated HG17, 18, 19 and 21, respectively) were isolated from the second library. A 6.6 kb SalI/EcoRI fragment of clone HG6 was found to hybridize with a 332 base pair HpaII/HpaII fragment of the mouse Egr-1 gene,

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which letter fragment spans the putative zinc finger region. The 6.6 kb fragment, in turn, was employed to 5 probe a cDNA library derived from human fibroblasts which have been stimulated for three hours with 20% fetal calf serum in the presence of 10  $\mu$ g/ml cyclohexamide. About 10,000 clones were screened and 10 the fifty positive clones obtained (designated "zap-1 through zap-50") are being subjected to nucleotide 15 sequence analysis. Preliminary sequence analysis 20 reveals that three clones, zap-2, zap-8, and zap-32, all encode the same transcript, namely a protein designated human EGR2, shown in Figure 3. Preliminary analysis 25 indicates approximately 92% homology between mouse Egr-1 and human EGR2 polypeptides in the zinc finger regions, but substantially less homology in the amino and carboxy terminal regions. Chromosome mapping studies, similar 30 to those described in Example 2, indicate that human chromosome 10, at bands q21-22, constitutes a locus for the human EGR2 gene.

30 The plasmid zap-32, containing the full length human EGR2 clone, was used as a probe in Southern blot analysis on DNAs from 58 unrelated Caucasians. It was 35 found that Hind III detects a simple two-allele polymorphism with bands at either 8.0 kb (A1) or 5.6 kb and 2.4 kb (A2). No constant bands were detected. The frequency of A1 was 0.90 and that of A2 was 0.10. No polymorphisms were detected for Apa I, BamH I, Ban II, Bgl I, Bgl II, BstE II, Dra I, EcoR I, EcoR V, Hinc II, Msp I, Pst I, Pvu II, Rsa I, Sac I, and Taq I in 10 unrelated individuals. Co-dominant segregation of the Hind III RFLP was observed in four large kindreds with a total of more than 350 individuals.

These data will be useful in gene linkage studies for mapping genes for certain genetic disorders. For example, the gene responsible for the dominantly inherited syndrome, multiple endocrine

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neoplasia, type 2A (MEN-2A) has been assigned by linkage to chromosome 10. Simpson, et al., Nature, 328, 528 (1987). Studies are currently underway to determine the linkage relationship between MEN-2A and EGR2 and are expected to be useful in cloning the MEN-2A gene as well as in serving as a diagnostic marker for the disease.

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EXAMPLE 7

15 Recombinant Expression Of Fusion Proteins

A 322 base HpaII/HpaII fragment (comprising nucleotides 1231-1553) derived from the OC3.1 cloned DNA was treated with DNA polymerase to fill in the single stranded ends. This fragment was inserted in plasmid pEX3 (obtained from K. Stanley, European Molecular Biology Laboratory, Postfach 10.2209, 6900 Heidelberg, F.R.G.) digested with SmaI. Stanley, K.K., et al., EMBO J., 3, 1429 (1984). This insertion placed the Egr-1 encoding DNA fragment in the same reading frame as plasmid DNA encoding cro- $\beta$ -galactosidase, allowing for the expression of a fusion protein comprising the amino terminal residues of cro- $\beta$ -galactosidase and 108 residues of Egr-1 amino acids 325 to 432. This cro- $\beta$ -galactosidase/Egr-1 fusion plasmid, designated pFIG, was used to transform E. coli NF1.

Induced (42°C) and un-induced (30°C) cultured cell lysates from growth of the transformed NF1 cells were then analyzed by SDS-PAGE. Upon Coomassie stain analysis, only induced cell lysates included an approximately 108 kd product, indicating presence of the projected expression product. Western blot analysis, using the rabbit polyclonal anti-peptide antibody VPS10 (see Example 4) raised against H-L-R-Q-K-D-K-K-A-D-K-S-C, confirmed that the fusion protein product contained Egr sequences.

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5        In a separate construction, a mouse Egr-1  
insert, from plasmid OC3.1, was fused, in frame, to a  
plasmid containing sequences from bovine growth hormone  
according to the methods described in Slamon, D.J., et  
10      al., Science, 233, 347 (1986). The resultant plasmid,  
designated pV4, comprised a fusion protein containing a  
fusion gene coding for bovine growth hormone amino acids  
1 to 192 and Egr-1 amino acids 2 to 533. This bGH/mouse  
15      Egr-1 DNA fusion plasmid, designated pV4, was expressed  
in E. coli and the resulting fusion protein, designated  
V4, was identified in Western blots by its reactivity  
with a bGH monoclonal antibody and its reactivity with  
20      VPS10 rabbit anti-Egr-1 peptide antiserum, prepared  
according to Example 4.

25      EXAMPLE 8

30      Determination of Egr  
Levels in Human Tumor and  
Non-Tumor Tissue

35      Using the mouse Egr-1 OC68 probe, Northern  
blot analyses were conducted to determine the levels of  
transcription of Egr protein encoding DNA in tumor  
versus surrounding normal tissue from resected human  
tumor specimens. The tumor samples were from lung (12),  
colon (7), colon mesastasis (1), bladder (1), rectal  
(1), giant cell (1), hepatoma (1), breast (1), MFH  
(malignant fibrous histiocytoma) (1), osteosarcoma (1)  
and rhabdomyosarcoma (1). In about 50% of these cases,  
there is markedly decreased (about three to ten-fold)  
expression of the Egr mRNA in tumor versus normal  
tissue. One implication of this finding is that Egr  
proteins of the invention may function as part of a  
negative regulatory pathway. In any event, it is clear  
that DNA sequences and antibodies of the invention are  
susceptible to use in differential diagnoses between  
tumorous and non-tumorous cell types.

It will be apparent from consideration of the foregoing illustrative examples that the present invention constitutes a substantial advance in the art and the achievement of a major goal in molecular biology, i.e., the characterization of genes which play a regulatory role in mammalian cell proliferation and differentiation. It will thus be understood that the information provided herein constitutes a basis for straightforward development of useful methods and materials not specifically the subject of the above examples. By way of illustration, possession of knowledge concerning the base sequence of cDNA and genomic DNA sequences encoding distinct mouse Egr-1 and human EGR2 early growth regulatory proteins comprising histidine-cysteine zinc finger amino acid sequences makes possible the isolation of other such structurally related proteins. The substantial homology between the zinc finger regions of Egr-1 and EGR2 coupled with lack of homology in other protein regions, when considered in light of the ability of Egr-1 probes to localize to human chromosome 5 while EGR2 probes localize to human chromosome 10, essentially assures the straightforward isolation of a human gene (provisionally designated "human EGRI") which encodes a protein more closely homologous to Egr-1 and a mouse gene (Egr-2) encoding a protein more closely homologous to EGR2.

While the above examples provide only limited illustration of in vitro and in vivo expression of DNA sequences of the invention, known recombinant techniques are readily applicable to development of a variety of prokaryotic and eucaryotic expression systems for the large scale production of Egr proteins and even development of gene therapy regimens.

Knowledge of the specifically illustrated mouse Egr-1 and human EGR2 proteins of the invention has been demonstrated to provide a basis for preparation of

highly useful antibodies, also provides a wealth of information concerning the nature of protein-nucleic acid interactions which, in turn, constitutes a basis for determination of significant early growth regulatory events. For example, and by analogy to steroid receptor protein structures, analysis of the structure of regions flanking the zinc fingers of Egr-1 and EGR2 and related proteins of the invention is expected to allow for identification of substances which may interact with the proteins to alter their DNA interactive capacities and thus provide the basis for inhibition or augmentation of their regulatory functions. Moreover, information available concerning specific events of DNA interaction of Egr proteins of the invention will permit, e.g., identification and use of potential competitive inhibitors of these proteins.

Just as Egr encoding DNA of the invention is conspicuously susceptible to use in differentiation of human tumor and non-tumor cells, antibodies prepared according to the invention are expected to be useful in differential screening of cells based on relative cellular concentrations of mRNA expression products and in the determination of specific genes susceptible to regulation by such products.

Because numerous modifications and variations in the practice of the present invention are expected to occur to those skilled in the art, only such limitations as appear in the appended claims should be placed thereon.

WHAT IS CLAIMED IS

5           1. A purified and isolated DNA sequence  
encoding a mammalian early growth regulatory protein  
which comprises one or more histidine-cysteine zinc  
10           finger amino acid sequences.

15           2. The DNA sequence according to claim 1  
encoding human EGR2 protein.

20           3. The DNA sequence according to claim 1  
encoding mouse Egr-1 protein.

25           4. The DNA sequence according to claim 1  
which is a cDNA sequence.

30           5. The DNA sequence according to claim 1  
which is a genomic DNA sequence.

35           6. The DNA sequence according to claim 1  
which is a wholly or partially synthetic DNA sequence.

35           7. The DNA sequence according to claim 1  
operatively associated with an homologous or  
heterologous expression control DNA sequence.

35           8. The DNA sequence according to claim 1  
selected from the group consisting of the DNA sequences  
set out in Figures 1A, 3, and 4.

35           9. A procaryotic or eucaryotic host cell  
transformed or transfected with a DNA sequence  
comprising a DNA sequence according to claim 1 or 7.

35           10. A viral or circular DNA plasmid vector  
comprising a DNA sequence according to claim 1.

11. The viral or circular DNA plasmid vector  
5 according to claim 10 further comprising an expression  
control DNA sequence operatively associated with said  
early growth regulatory protein encoding DNA.

10 12. A method for the production of an early  
growth regulatory protein comprising:  
15 growing, in culture, a host cell transformed  
or transfected with a DNA sequence according to claim 1;  
and  
20 isolating from said host cell or culture the  
polypeptide product of the expression of said DNA  
sequence.

25 13. A method for the production of an early  
growth regulatory protein comprising:  
disposing a DNA sequence according to claim 1  
in a cell free transcription and translation system; and  
30 isolating from said system the polypeptide  
product of the expression of said DNA sequence.

35 14. The polypeptide product of the in vitro  
or in vivo expression of part or all of a protein encoding  
region of a DNA sequence according to claim 1.

15. The polypeptide product according to  
claim 14 which is a fusion protein comprising part or  
all of a mammalian early growth regulatory protein which  
comprises one or more histidine-cysteine zinc finger  
amino acid sequences and part or all of a heterologous  
proteins.

16. The polypeptide product according to  
claim 15, which comprises a fusion of cro- $\beta$ -  
galactosidase and Egr-1 amino acid sequences or bovine  
growth hormone and Egr-1 amino acid sequences.

17. A purified and isolated mammalian early  
5 growth regulatory protein comprising one or more histi-  
dine-cysteine zinc finger amino acid sequences.

18. The protein according to claim 17 and  
10 having the amino acid sequence set out in Figure 1A.

19. The protein according to claim 17 and  
15 having the amino acid sequence set out in Figure 3.

20. A synthetic peptide duplicative of a  
sequence of amino acids present in a mammalian early  
growth regulatory protein comprising one or more histi-  
dine-cysteine zinc finger amino acid sequences and  
25 sharing at least one antigenic epitope of such  
protein.

21. The synthetic peptide of claim 20 and  
30 having an amino acid sequence partially duplicative of  
the amino acid sequence set out in Figure 1A.

22. The synthetic peptide of claim 20 and  
35 having an amino acid sequence partially duplicative of  
the amino acid sequence set out in Figure 3.

23. The synthetic peptide of claim 20 and  
further characterized as duplicative of an amino acid  
sequence not involved in DNA binding functions.

24. An antibody specifically immunoreactive  
with at least one epitope of a mammalian early growth  
regulatory protein comprising one or more histidine-  
cysteine zinc finger amino acid sequences.

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25. The antibody according to claim 24  
wherein said epitope is other than an epitope within the  
5 DNA binding functional region thereof.

10 26. The antibody according to claim 24  
selected from the group consisting of monoclonal and  
polyclonal antibodies.

15 27. The antibody according to claim 24  
capable of specifically binding with a proteinaceous  
material comprising an amino acid sequence duplicating  
an antigenic epitope within the following amino acid  
20 sequence:

H-L-R-Q-K-D-K-K-A-D-K-S-C.

25 28. The antibody according to claim 24  
capable of specifically binding with a proteinaceous  
material comprising an amino acid sequence duplicating  
an antigenic epitope within the following amino acid  
30 sequence:

C-G-R-K-F-A-R-S-D-E-R-K-R-H-T-K-I.

35 29. A method for quantitative detection of a  
mammalian early growth regulatory protein comprising one  
or more histidine-cysteine zinc finger amino acid  
sequences based on the immunological reaction of the  
same with an antibody according to claim 24.

30. A method for quantitative detection  
within a sample of messenger RNA transcripts for  
mammalian early growth regulatory proteins comprising  
one or more histidine-cysteine zinc finger amino acid  
sequences comprising the step of hybridizing RNA within  
said sample with a DNA sequence partially or wholly  
duplicating a DNA sequence according to claim 1.

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31. A method for quantitative detection  
within a sample of DNA encoding for mammalian early  
5 growth regulatory proteins comprising one or more  
histidine-cysteine zinc finger amino acid sequences  
comprising the step of hybridizing DNA within said  
sample with a DNA sequence partially or wholly  
10 duplicating a DNA sequence according to claim 1.

32. A method for detecting a disease  
15 genetically linked to a mammalian Egr gene comprising  
the step of quantitating mammalian early growth  
regulatory DNA sequences according to claim 1.

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10      20      30      40      50      60  
GGGGAGCCGGCCGGATTCGGCCGCCGGCCAGCTTCCGGCCGGCAAGATCGGGCC

70      80      90      100     110     120  
TGCCCCAGCCTCCGGCGGAGCCCTGCGCTCCACCACGGCCGGCTACCGCCAGCCTGGG

130     140     150     160     170     180  
GGCCCCACCTACACTCCCCGGCAGTGTGCCCTGGCACCCGGCATGTAAACCCGGCCAAACCCCC

190     200     210     220     230     240  
GGCGAGTGTGCCCTCAGTAGCTTTCGGCCGGCTGGCCACCCAAACATCAGTTCT

250     260     270     280     290     300  
CCAGCTCGCTGGCTGGGGATGGCAGGCCAAGGCCGGAGATGCCAATTGATGCTCTCGCTG  
MetAlaAlaAlaLysAlaGluMetGlnLeuMetSerProLeu

310     320     330     340     350     360  
CAGATCTGACCCGGTTCGGCTCCTCACTCACCCACCATGACAACCTACCCCAA  
GlnIleSerAspProPheGlySerPheProHisSerProThrMetAspAsnTyrProLys

370     380     390     400     410     420  
CTGGAGGAGATGCTGAGCAACGGGGCTCCCGCAGTTCCTCGGTGCTGCCGGAAACC  
LeuGluGluMetMetLeuLeuSerAsnGlyAlaProGlnPheLeuGlyAlaAlaGlyThr

FIGURE 1.1

430            440            450            460            470            480  
 CCAGAGGGCAGGGGGCTAAATTAGCAGCACCAGCAGGAAACCCGAGGCGGGTGGGGGG  
 ProGluGlySerGlyGlyAsnSerSerSerSerSerSerSerSerSerSerSerSerSer  
  
 490            500            510            520            530            540  
 GGCAGCAAACAGGGCAGGGCCTTCATTCTCAAGGGAGCCGAGCGAACAAACCCCTAT  
 GlySerAsnSerGlySerSerAlaPheAsnProGlnGlyGluProSerGluGlnProTyr  
  
 550            560            570            580            590            600  
 GAGCACCTGACACAGAGTCTTCTGACATCGCTCTGAATAATGAGAAGGGCGATGGTC  
 GluHisLeuThrThrGluSerPheSerAspIleAlaLeuAsnAsnGluLysAlaMetVal  
  
 610            620            630            640            650            660  
 GAGACGAGTTATCCCAGCCAAACGACTCTCGGTTGCCATCACCTATACTGCCGCTTC  
 GluThrSerTyrProSerGlnThrThrArgLeuProProProIleThrTyrThrGlyArgPhe  
  
 670            680            690            700            710            720  
 TCCCTGGAGCCGGCACCCAAACAGTGGCAACACACTTGTGGCCTGAACCCCTTTCAAGCCTA  
 SerLeuGluProAlaProAsnSerGlyAsnThrLeuTrpProGluProLeuPheSerLeu  
  
 730            740            750            760            770            780  
 GTCAGTGGCTCGTGAGGCATGACCAAATCCTCCGACCTCTTCATCCATCGGGCCTTCTCCA  
 ValSerGlyLeuValSerMetThrAsnProProThrSerSerSerAlaProSerPro  
  
 790            800            810            820            830            840  
 GCTGCTTCATCGTCTCCTCTGCCTCCAGAGCCCCCCCCTGAGCTGTGCCGCGTCC  
 AlaAlaSerSerSerSerSerAlaSerGlnSerProProLeuSerCysAlaValProSer

FIGURE 1.2

850	860	870	880	890	900
AACGACAGCAGTCCC	CATCTACTCGGCTGCC	CACCTTCTACTCCCC	AAACACTGACAT		
AsnAspSerSerProIle	TerProIleTerSerAla	AlaProThrProThr	ProAsnThrAspIle		
910	920	930	940	950	960
TTTCCTGAGCCCCAAAGCC	AGGCCCTTCC	GGCTCGG	CAGGCACAGCCTT	GCAGTACCCG	
PheProGluProGlnSerGln	AlaPhe	ProGlySerAla	GlySerAla	GlySerAla	TerPro
970	980	990	1000	1010	1020
CCTCCTGCCTACCC	TGCCAACCAAGGT	TCCAGGT	TCCATGATCCC	CTGACTATCTG	
ProProAlaTerProAla	TerIleTerIle	TerIleTerIle	TerIleTerIle	TerIleTerIle	
1030	1040	1050	1060	1070	1080
TTTCCACAAACAGGGAGACCTGAGC	ACCTGGC	CACCCAGACCAG	ACCCAGAACG	CCCTTCCAGGGT	
PheProGlnGlnGlyAsp	LeuSerLeuGly	TerGlyGly	TerGlyGly	TerGlyGly	
1090	1100	1110	1120	1130	1140
CTGGAGAACCGTACCC	CAGCAGC	ACTCCACTATCC	ACTATCCACT	ATTAAAGCC	TTCGCC
LeuGluAsnArgTerGln	TerGln	TerGln	TerGln	TerGln	TerGln
1150	1160	1170	1180	1190	1200
ACTCAGTCGGGCTCCC	CAGGACTTAAGGCT	CTTAATACCACCT	ACCAATCCC	CAGCTC	CATC
ThrGlnSerGlySerGln	TerGlySerGln	TerGlySerGln	TerGlySerGln	TerGlySerGln	TerGlySerGln
1210	1220	1230	1240	1250	1260
AAACCCAGCCGCATGGC	AAAGTACCC	AAACCGGCCC	AAACCGGACAC	CCCCCATGAA	ACGCG
LysProSerArgMetArg	TerLysTer	TerLysTer	TerLysTer	TerLysTer	TerLysTer

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1270      1280      1290      1300      1310      1320  
 CCATATGCTTGCCTGTGAGTCCTGCGATTCGCCGCTTCTCGATGAGGCTTAC  
 ProTyrAlaCysProValGlusercysAspArgSerArgSerAspGluLeuThr  
 1330      1340      1350      1360      1370      1380  
 CGCCATATCCGCATCCACAGGCCAGAAGCCCTTCCAGTGTGAATCTGCATGGCTAAC  
 ArgHisIleArgIleHisthrGlyGlnLysProPheGlnCysArgIleCysMetArgAsn

1390      1400      1410      1420      1430      1440  
 TTCACTCGTAGTGAACCACTTACCAACATCCGCACACACAGGCCAGAAGGCCTTT  
 PheSerArgSerAspHisIleLeuThrThrHisIleArgThrHisthrGlyGluLysProPhe

1450      1460      1470      1480      1490      1500  
 GCCTGTGACATTGTGGAGGAAGTTGCCAGGAGTGTGAACGCAAGAGGCATAACAAA  
 AlaCysAspIleCysGlyArgLysPheAlaArgSerAspGluArgLysArgHisthrLys

1510      1520      1530      1540      1550      1560  
 ATCCATTAGACAGAACAGAACAAAGCAGACAAAGAACAAAGTGTGGCTCCCCGGCTGCC  
 IleHisIleLeuArgGlnLysAspLysLysAlaAspLysSerValValAlaSerProAlaAla

1570      1580      1590      1600      1610      1620  
 TCTTCACTCTTACCCATCCCCAGTGGCTACCTCCTACCACTCCCTGCACCA  
 SerSerLeuSerSerTyrProSerProValAlaThrSerTyrProSerProAlaThrThr

1630      1640      1650      1660      1670      1680  
 TCATTCCCATCCCCCTGTGCCCACTTCCCTACTCCCTACTCCCTGCCTCCACCTACCCATCT  
 SerPheProSerProValProThrSerSerTyrSerSerProGlySerSerThrTyrProSer

FIGURE 1.4

1690 1700 1710 1720 1730 1740  
CCTGGCACAGTGGCTTCCCGTCAGTGGCCACCTTGCCTTCCGTTCAGCTTCCACT  
ProAlaHisSerGlyPheProSerProSerValAlaThrThrPheAlaSerValProPro

1750 1760 1770 1780 1790 1800  
GCTTTCCCCACCCAGGTCAAGCAGCTTCCCGTCTGCAGGCGTCAGCAGCTTCCAGCACC  
AlaPheProThrGlnValSerSerPheProSerAlaGlyValSerSerPheSerThr  
1810 1820 1830 1840 1850 1860  
TCAACTGGTCTTTCAAGACATGACAGCGACCTTCTCCAGGACAAATTGAAATTGCTAA  
SerThrGlyLeuSerAspMetThrAlaThrPheSerProArgThrIleGluIleCys

1870 1880 1890 1900 1910 1920  
AGGGATAAAAGCAAAGGGAGGGCAGGAAGACATAAAGGCACAGGAGGGAAAGAG

1930 1940 1950 1960 1970 1980  
ATGGCCGCAAGAGGGGCCACCTCTAGGTCAAGATGGAAAGATCTCAAGGCCAAGTCCTTCT

1990 2000 2010 2020 2030 2040  
ACTCACGAGTAGAAGGGACCCGTTGGCCAACAGGCCCTTCACCTTACCATCCCTGCCTCCCC

2050 2060 2070 2080 2090 2100  
GTCCCTGTTCCCTTCACTTCAGCTGGCTGAAACAGGCCATGTCCTCACCTCTAT

FIGURE 1.5

2110 2120 2130 2140 2150 2160  
CAAAGGACTTGATTGCATGGTATGGATAAATCATTTCAAGTATCCTCCATCACATG

2170 2180 2190 2200 2210 2220  
CCTGGCCCTTGCCTTCAGGCCTAGACCATCAAGTGGCATAAAGAAAAAAATGGG

2230 2240 2250 2260 2270 2280  
TTTGGCCCTCAGAACCCCTGCCCTGCATCTTGTACAGGCATCTGTGCCATGGATTGT  
2290 2300 2310 2320 2330 2340  
TTCCTGGGTATTCTTGATGTGAAGGATAATTGCATACTCTATTGTATTGGAGT

2350 2360 2370 2380 2390 2400  
AAATCCCTCACTTTGGGGAGGGGGAGCAAAGCAAACCAATTGATGATCCCTCTATT

2410 2420 2430 2440 2450 2460  
TTGTGATGACTCTGCTGTGACATTAGGTTTGAAGGCATTTTTTCAAGGCAGGTCCCT

2470 2480 2490 2500 2510 2520  
AGGTATTAACTGGGACCATGTTGTCAGAGTGTGTTCCGTTAATTGTAAATACTGGCTCG

FIGURE 1.6

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2530 2540 2550 2560 2570 2580  
ACTGTAACCTCACATGTGACAAAGTATGTTGGTTGGCTTTGAGAAT

2590 2600 2610 2620 2630 2640  
TTTTTGGCCGGTCCCTTTGGTTCAAAAGTTTCAACGTCTTGCTGCCTTTGCTGACACG

2650 2660 2670 2680 2690 2700  
CCTTCGGATGGCTTGCACATGGCAGATGTTGGGACACGGCTCACCTTACGCCTTAAGGGGG  
2710 2720 2730 2740 2750 2760  
TAGGAGTGTGTTGGGGAGGGCTTGAGAGCAAAACGAGGAAAGAGGGCTGAGCTGAGC

2770 2780 2790 2800 2810 2820  
TTTCGGTCTCAGAAATGTAAGAAGAAATTTAAACAAAATCTGAACCTCTCAAAAGTC

FIGURE 1.7

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2830 2840 2850 2860 2870 2880  
TATTTTCTAAACTGAAATGTAATTATACATCTATTCAAGGAGTTCGGAGTGTGTGGT

2890 2900 2910 2920 2930 2940  
TACCTACTGAGTAGGGCTGCAGTTTGTATGTTATGAACATGAAGTCATTATTTGTGG

2950 2960 2970 2980 2990 3000  
TTTTTACTTTGTACTTGTGTGTTGCTTAACCAAGTAACCTGTTGGCTTATAACA

3010 3020 3030 3040 3050 3060  
CATTGAATGGCGCTCTATTGCCCATGGGATATGTGGTGTATCCTTCAGAAAAATTAAAA

3070 3080  
GGAAAAATAAAAAAA

SUBSTITUTE SHEET

FIGURE 1.8

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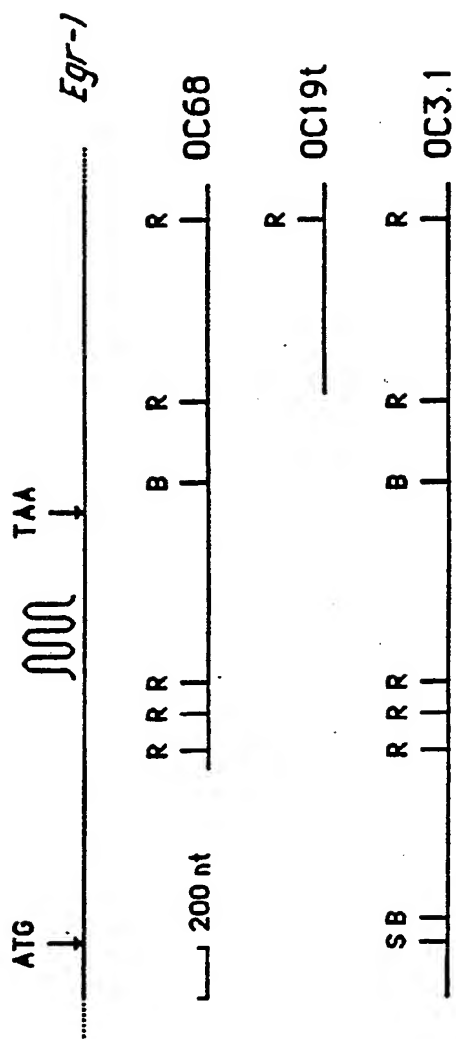


FIGURE 1B

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"zinc finger"  
consensus sequence

Murine Egr-1

P H E R P Y A C P V E S C D R R F S R S D E L T R H I R I H  
T G Q K P F Q C - - R I C M R N F S R S D H L T T H I R T H  
T G E K P F A C - - D I C G R K F A R S D E R K R H T K I H

Drosophila Kruppel

S R D K S F T C K I - - C S R S F G Y K H V L Q N H E R T H  
T G E K P F E C P E - - C D K R F T R D H H L K T H M R L H  
T G E K P Y H C S H - - C D R Q F V Q V A N L R R H L R V H  
T G E R P Y T C E I - - C D G K F S D S N Q L K S H M L V H

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Xenopus TFIIIA  
(finger 2)

T G E K P F P C K E E G C E K G F T S L H H L T R H S L T H

FIGURE 2

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10 20 30 40 50 60  
TTTTTTTTGGTGTGTGGTTAAAGTGTGGAGGGCAAAAGGAGATACCA

70 80 90 100 110 120  
TCCCCAGGCTCAGTCCAACCCCTCTCCAAAACNGTGTCTGACACTCCAGGTAGCGA

130 140 150 160 170 180  
GGGACTGGGTCTCAGGTCTGGAGGAGCAAAATGATGACCCCCAAGGCCGTAGACAAA  
MetMetThrAlaLysAlaValAlaAspLys

190 200 210 220 230 240  
ATCCCAGTAACCTCTCAGTGTGTGTGGCACCCAGCTGTGACAACATCTACCCGGTGGAG  
IleProValThrLeuSerGlyPheValHisGlnLeuSerAspAsnIleTyrProValGlu

250 260 270 280 290 300  
GACCTCGCCGCCACGTGGTGAACCATCTTCCCCAATGCCGAACATGGGAGGGCCCTTGTAC  
AspLeuAlaAlaThrSerValThrIlePheProAsnAlaGluLeuGlyGlyProPheAsp

310 320 330 340 350 360  
CAGATGAACGGAGTGGCCGGAGATGGCATGATCAACATGACATGACTGGAGAGAAGAGG  
GlnMetAsnGlyValAlaGlyAspGlyMetIleAsnIleAspMetThrGlyGluLysArg

FIGURE 3.1

SUBSTITUTE SHEET

370	380	390	400	410	420
TCGTTGGATCTCCCATATCCCAGCAGCTTGTCTCCGTCTGCACCTAGAAACCGAGACC					
SerLeuAspLeuProTyrProSerSerPheAlaProValSerAlaProArgAsnGlnThr					
430	440	450	460	470	480
TTCACTTACATGGCCAACTTCTCCATTGACCCACAGTACCCACTGGTGGCCAGCTGCTACCCA					
PheThrTyrMetGlyLysPheSerIleAspProGlnTyrProGlyAlaSerCystYrPro					
490	500	510	520	530	540
GAAGGCATAATTCAATTATTGTGAGTGCAGGCATCTGGCAAGGGTCACTTCCCAAGCTTCA					
GluGlyIleIleLeuAsnIleValSerAlaGlyIleLeuGlnGlyValThrSerProAlaSer					
550	560	570	580	590	600
ACCACAGCCTCATCCAGCGTCAACCTCTGGCTTCCCAACCCACTGGCCACAGGACCCCTG					
ThrThrAlaSerSerValThrSerAlaSerProAlaSerProAsnProLeuAlaThrGlyProLeu					
610	620	630	640	650	660
GGTGTGGCACCATGTCAGCCAGCCAGCCTGACCTGGACCCACCTGTACTCTCCGCCACCG					
GlyValCysThrMetSerGlnThrGlnProAspLeuAspHisLeuTyrSerProProPro					
670	680	690	700	710	720
CCTCCCTCCTTATTCTGGCTCTGGAGAGACCTTACCAAGAACCCCTCTGGCGTTCCCTG					
ProProProTyrSerGlyCysAlaGlyAspIleuTyrGlnAspProSerAlaPheLeu					

## SUBSTITUTE SHEET

## SUBSTITUTE SHEET

FIGURE 3.3

1090 1100 1110 1120 1130 1140  
TATAACCACCCACTGCCACTGGCCATTCTGAGGCCCTCGCAAGTACCCCAACAGA  
TyrAsnProHisHisLeuProLeuArgProIleLeuArgProArgLysTyrProAsnArg

1150 1160 1170 1180 1190 1200  
CCAGCAAGACGCCGGTGCACGAGAGGGCCCTACCCGTCAGCAGAAGGGCTGCGACCGG  
ProSerLysThrProValHisGluArgProTyrProCysProAlaGluGlyCysAspArg

1210 1220 1230 1240 1250 1260  
CGGTCTCCGGCTCTGACGAGCTGACACGGCACATCCGAATCCACACTGGCATAAGCCC  
ArgPheSerArgSerAspGluLeuThrArgHisIleArgIleHisthrglyHisLysPro

1270 1280 1290 1300 1310 1320  
TTCCAGTGCGATCTGGCATGGCAAACCTCAGCCGCAGTGACCACTCACCCATATC  
PheGlnCysArgIleCysMetArgAsnPheSerArgSerAspHisLeuThrThrHisIle

1330 1340 1350 1360 1370 1380  
CGCACCCACACCGGTGAGAACGCCCTCGCCTGTGACTACTGTGCCGAAAGTTGCCCGG  
ArgThrHisThrGlyGluLysProPheAlaCysAspTyrCysGlyArgLysPheAlaArg

1390 1400 1410 1420 1430 1440  
AGTGATGAGGAAACGCCACCAAGATCCACCTGAGACAGAAAGAGCGGAAAGGCAGT  
SerAspGluArgLysArgHisIleHisIleLysGluArgGlnLysGluArgLysSerSer

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1450 1460 1470 1480 1490 1500  
GCCGCCCTCTGCATCGGTGCCAGCCCCCTCTACAGCCTCCTGCTCTGGGGCGTGCAGGCC  
AlaProSerAlaSerValProAlaProSerThrAlaSerCysSerGlyGlyValGlnAla

1510 1520 1530 1540 1550 1560  
TGGGGGTACCCCTCTGCAGGCAAGTAACAGCAGGCAAGTCTTGAGGGAGGGGGCTTG  
TrpGlyTyrProValGlnEnd

1570 1580 1590 1600 1610 1620  
CTCCCTCTCGGACCCGGACACCTTGAGATGAGACTCAGGCTGATAACACCAGCTCCAAAGG

1630 1640 1650 1660 1670 1680  
TCCCGGGCCCTTTCTCCACTGGAGCTGGCACAAACAAACACTACCACCCCTTTCCTGTCCC

1690 1700 1710 1720 1730 1740  
TCTCTCCCTTTCTGGCAAAAGGCTTGGGGAGCTTAGGCACCTGCCCTTTCACCTAG

1750 1760 1770 1780 1790 1800  
AAGCAGGGTTCTTCTAAACTAGCCCCATTAGCTCTAGTCCTCTAGGTGAGTTGACTATCAA

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FIGURE 3.5

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1810	1820	1830	1840	1850	1860
CCCAAGGCCAAAGGGGAGGCTCAGAAGGAGGTGGTGTGGGATCCCCCTGGCCAAGAGGGCT					
1870	1880	1890	1900	1910	1920
GAGGCTCTGACCCCTGCTTAAAGGGCTTGTGACTTAGGTTTTGCTACCCCACTTCCCTTA					
1930	1940	1950	1960	1970	1980
TTTGACCCATCACAGGTTTTGACCCCTGGATGTCAGAGTTGATCTAAGACGGTTTCTAC					
1990	2000	2010	2020	2030	2040
AATAGGTTGGGAGATGCTGATCCCTCAAGTGGGACAGCCAAAAGACAAGCAAACCTGA					
2050	2060	2070	2080	2090	2100
TGTGCACTTATGGCTGGGACTGATTGGGACATTTGACACTGAGTGAAAGTATAAGCC					
2110	2120	2130	2140	2150	2160
TTTATGCCACACTCTGTGGCCCTAAATGGTGAATCAGAGCATATCTAGTTGTCTCAACC					

FIGURE 3.6

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2170 2180 2190 2200 2210 2220  
CTTGAAGCAATATGTATTATACTCAGAGAACAGAACAGAACAGAACAGAACGGT

2230 2240 2250 2260 2270 2280  
AGCAATATCTGCTCCCTTTCGAGTTCTTCTAGAAATTGTTAGGCTATTTCAGTGTATAT

2290 2300 2310 2320 2330 2340  
CCACTCAGATTTGTGTATTGTACACTGTACACACTGTCTCTAAATTCTGAATTCTTGT

2350 2360 2370 2380 2390 2400  
GGAAAAATGTAAGCATTATGATCTCAGAGGTAACTTAACTTAAGGGGATGTACATA

2410 2420 2430 2440 2450 2460  
TTCTCTGAAACTTAGGATGCATTGCAATTGTTGGAAGTGTCCCTGGCTCGCCTTGTGTGAT

2470 2480 2490 2500 2510 2520  
GTAGACAAATGTTACAAGGCTGCATGTAATTGGTTGCCTTATGGAGAAAAAAATCA

SUBSTITUTE for FIGURE

FIGURE 3.7

2530 2540 2550 2560 2570 2580  
CTCCCTGACTTTAGTATGGCTGTATATTGCTATTAAATTCAAAATTTCATTTTTTTAG

2590 2600 2610 2620 2630 2640  
AGTATATTGTTGCTATGCTTGTGACTTTAAAGTGTACCTTGTAGTCAAATTTC

2650 2660 2670 2680 2690 2700  
AGATAAGAACATGACATAATTGTTACCGGAGCTGANNNNTGGTTGGTCATTAGCTCTTAATA

2710 2720 2730 2740 2750 2760  
GTTGTAACAAATAATCTATTCTAACGGAAAACCAACTAACGAACTGAAGTTCAAGATAATGG

2770 2780 2790 2800 2810  
ATGGTTGTGACTTATAGTGTAAATAACTTTCAACAAAAAA

FIGURE 3.8

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**SUBSTITUTE SHEET**

FIGURE 4.1

-335 ATAATGGCCC TGCCGCTTCC GGTCTGGAA GGAAGGGCGAA GCGGGGGTTC GGGGGGGG  
 -326

-315 AGGCTGGAA CTCCAGGGCC CTGGGGGGGG AGGGCCACTGC TGTGTCCA ATACTGGG  
 -316 <sup>Sma</sup>I

-315 TTCCAGGGAC CTGAGGGCTC GCGATGCGGG AGCGGGGTGCG AGGGTGGGG TGCCACAC  
 -316

-315 TCTGGGGATG GAGGGCTCA CGTCACTCCG GGTCTCCCG AGCGGGCTT CGATATTAGG  
 -316

-315 GCTTCCTCT TCCCCATATAT GGGCATATAC GTCACGGCG AGGGGGGGG GTGCTGTCC  
 -316

-315 AGACCCCTGA ATAAGGGCC GATTGGGGAA GTCGGCGAGG ATCCACGGC GCGAACTT  
 -316

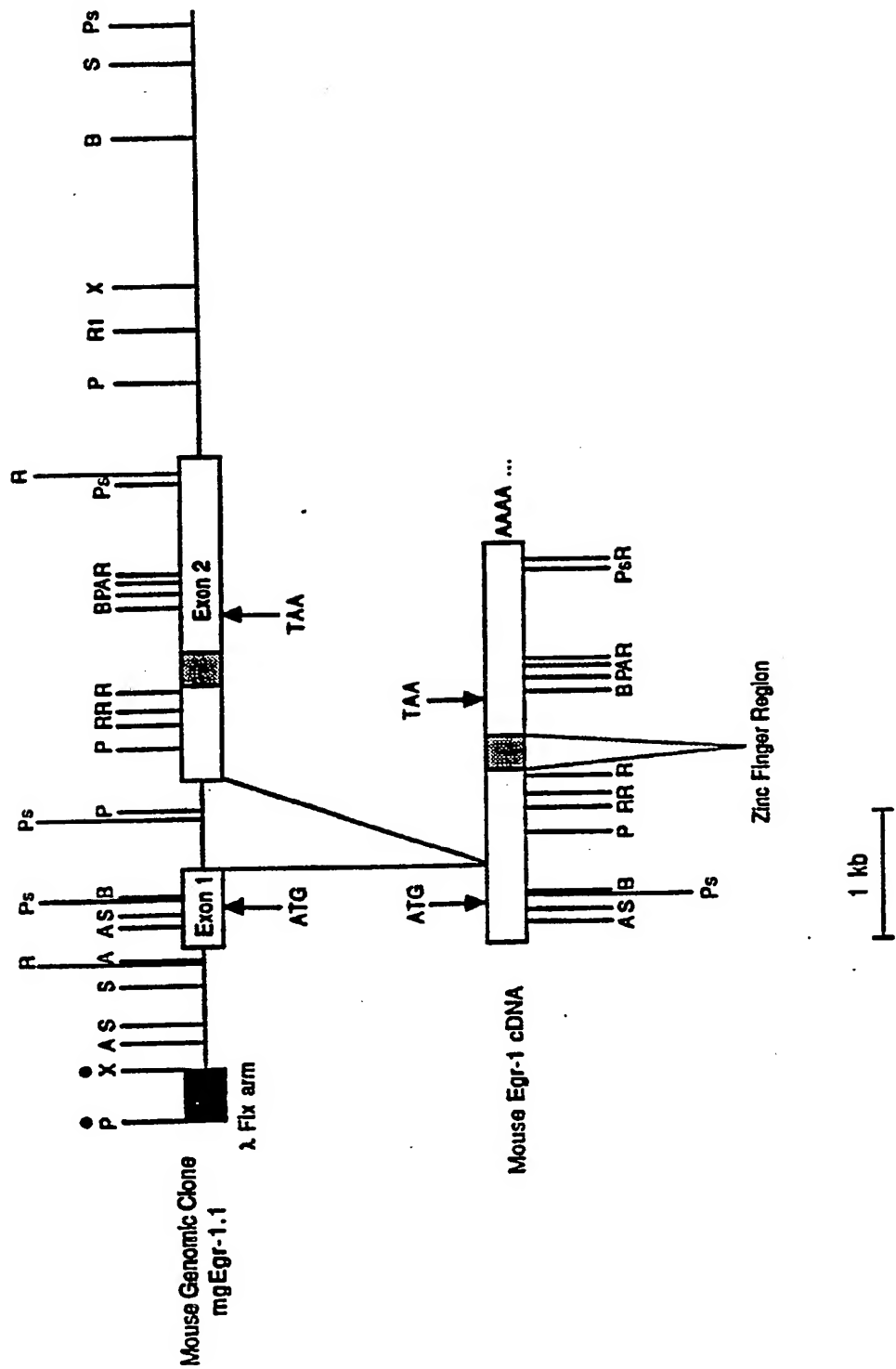
-316 GGGAAACGCC GTCGGGATTC GGCACCGCGG CGAGCTTCCG CGGGGGAAAG ATCGGGCCCT  
 -316 <sup>Sma</sup>I

+86 GCGCCAGCTC CGGGCGAGC CCTGGTCCA CGACGGGGCG CGGTACCCGC CAGGCTGGGG  
 +116 GCGCACCTAC ACTCCCGCA GTGTCGGCT GCAACCCCGG CGAACCCCGG  
 +116

+126 GCGAGTGTTC CCTCAATAGC TTGGGGGGGG GGCTGGGGCC ACCACCCAAAC ATCAAGTTC  
 +126 <sup>Sma</sup>I

FIGURE 4.2

SUBSTITUTE SHEET



**SUBSTITUTE SHEET**

# INTERNATIONAL SEARCH REPORT

International Application No. PCT/US89/01473

## I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) <sup>6</sup>

According to International Patent Classification (IPC) or to both National Classification and IPC  
 C07H 15/12, C12P 21/00, 19/34; C12N 15/00, 7/00, 1/20;  
 C12Q 1/68; C07K 13/00

## II. FIELDS SEARCHED

Minimum Documentation Searched <sup>7</sup>

Classification System	Classification Symbols
US	536/27; 435/68, 91, 252.3, 6,530/350, 387      172.1, 172.3, 320,

Documentation Searched other than Minimum Documentation  
to the Extent that such Documents are Included in the Fields Searched <sup>8</sup>

**CAS database (1967-1989) keywords: finger/protein/motif  
early growth regulatory/gene/expression/sequence**

## III. DOCUMENTS CONSIDERED TO BE RELEVANT <sup>9</sup>

Category <sup>10</sup>	Citation of Document, <sup>11</sup> with indication, where appropriate, of the relevant passages <sup>12</sup>	Relevant to Claim No. <sup>13</sup>
X Y	Chavrier, P. et al (January 1988) EMBO Journal, Volume 7: 29-35; "A gene encoding a protein with zinc fingers is activated during G <sub>0</sub> /G <sub>1</sub> transition in cultured cells. See entire document.	1-5, 8-10 6, 7, 11, 12 14-16
X Y	Chowdhury, K. et al (March 1988) Cell, Volume 48: 771-778; "A Multigene family encoding several finger structures is present and differentially active in mammalian genomes". See entire document.	1-5, 8-10 6, 7, 11, 12, 14-16
X Y	Lau, L. et al (March 1987) PNAS, Volume 84: 1182-1186; "Expression of a set of growth-related immediate early genes in BALB/c 3T3 cells: coordinate regulation with c-fos or c-myc. See entire document	1-5, 7-11 6, 12, 14-16

\* Special categories of cited documents: <sup>10</sup>

"A" document defining the general state of the art which is not considered to be of particular relevance

"E" earlier document but published on or after the international filing date

"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

"O" document referring to an oral-disclosure, use, exhibition or other means

"P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"A" document member of the same patent family

## IV. CERTIFICATION

Date of the Actual Completion of the International Search

Date of Mailing of this International Search Report

27 June 1989

24 AUG 1989

International Searching Authority

Signature of Authorized Officer

ISA US

*Anne Brown*  
Anne Brown

## FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET

P,Y	<p>Pannuti, A. et al (May 1988) Nucleic Acids Res., Volume 16: 4227-4237 "Isolation of cDNAs encoding finger proteins and measurement of the corresponding mRNA levels during myeloid terminal differentiation" See entire document.</p> <p>Almendrol et al (May 1988) Mol Cell Biol, Vol. 8:2140-2148 "Complexity of the Early Genetic Response to Growth Factors in Mouse Fibroblasts". See entire document.</p>	1-12, 14-16 1-12, 14-16
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V.  OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE<sup>1</sup>

This International search report has not been established in respect of certain claims under Article 17(2) (a) for the following reasons:

1.  Claim numbers \_\_\_\_\_, because they relate to subject matter<sup>1,2</sup> not required to be searched by this Authority, namely:

2.  Claim numbers \_\_\_\_\_, because they relate to parts of the International application that do not comply with the prescribed requirements to such an extent that no meaningful International search can be carried out<sup>1,2</sup>, specifically:

3.  Claim numbers \_\_\_\_\_, because they are dependent claims not drafted in accordance with the second and third sentences of PCT Rule 6.4(a).

VI.  OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING<sup>2</sup>

This International Searching Authority found multiple inventions in this International application as follows:

## See attached sheet

1.  As all required additional search fees were timely paid by the applicant, this International search report covers all searchable claims of the International application.

2.  As only some of the required additional search fees were timely paid by the applicant, this International search report covers only those claims of the International application for which fees were paid, specifically claims:

3.  No required additional search fees were timely paid by the applicant. Consequently, this International search report is restricted to the invention first mentioned in the claims; it is covered by claim numbers: 1-12, 14-16

4.  As all searchable claims could be searched without effort justifying an additional fee, the International Searching Authority did not invite payment of any additional fee.

## Remark on Protest

The additional search fees were accompanied by applicant's protest.  
 No protest accompanied the payment of additional search fees.

ATTACHMENT TO PCT/ISA/210

VI. OBSERVATION WHERE UNITY OF INVENTION IS LACKING

I. Claims 1-12, and 14-16 are drawn to EGR DNA sequence, vector or cell containing it, transformed host, method of using vector or cell and a polypeptide.

II. Claim 13 is drawn to cell-free method of preparing an early growth regulatory protein.

III. Claims 17-19 are drawn to native EGR protein.

IV. Claims 20-23 are drawn to synthetic peptide fragment antigenically related to and containing homology to native EGR protein.

V. Claims 24-29 are drawn to antibodies to EGR, anti-bodies to region EGR, method of using antibody.

VI. Claim 30 is drawn to method of detecting mRNA.

VII. Claim 31 is drawn to method of detecting DNA.

VIII. Claim 32 is drawn to method of diagnosis.

The claims do not embrace one single general inventive concept as defined in Rule 13.

Groups I, III, IV and V contain claims directed to distinct chemical entities. These are a gene and a protein coded for by that gene, a native EGR protein, a synthetic peptide fragment, antibodies which bind to EGR proteins. Groups II, VI, VII and VIII are directed to alternative uses for the nucleotide defined in Group I.

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